

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of	:	
Dick, Stefan O., et al.	:	
	:	Art Unit:
Serial No.	:	
	:	Examiner:
Filing Date	:	
	:	
Attorney Docket No. P-1047	:	
	:	
For: PLATE-SHAPED PRESSED BODIES	:	

Box Patent Application
Assistant Commissioner for Patents
Washington, D.C. 20231

PRELIMINARY AMENDMENT

In the Specification

A provisional application Serial No. 60/260,282 was filed on January 8, 2001. The attached non-provisional application claims priority under 35 U.S.C. Section 119(e) from this earlier filed provisional application. The provisional application was the translation of a German patent application. The translation contained a number of awkward phrases and was not in proper form for review by the United States Patent and Trademark Office. For example, the one figure was not specifically referred to in a separate section of the application. In addition, a substantial number of minor changes were made to the text of the application to put it in proper condition for review by the United States Patent

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and Trademark Office. Attached as Exhibit A is a copy of the provisional application with the modifications made to that application noted on the pages in handwriting. A clean copy of that application is the original, non-provisional application that is being filed with the United States Patent and Trademark Office on this date. No new subject matter is introduced by any of the amendments to the specification.

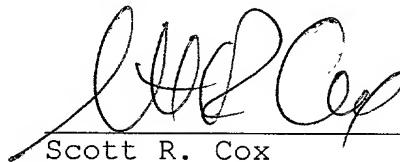
In the Claims

All of the claims that were filed with the provisional application are cancelled and replaced by new Claims 1-24, which are included with the attached non-provisional application.

CONCLUSION

The applicants believe that the application is in condition for review by the United States Patent and Trademark Office. If there are any questions, please contact applicants' counsel.

Respectfully submitted,



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Attachment

CERTIFICATE OF SERVICE

I hereby certify that this correspondence is being deposited with the United States Postal Service in an envelope as Express Mail Post Office to Addressee," mailing Label Number ET998036205US, addressed to Box Patent Application, Assistant Commissioner for Patents, Washington, D.C. 20231.

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Dorothy Goodlett

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~~December 21, 2000~~

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~~Translation from German~~

Title

Plate-shaped pressed bodies

Field of Invention Description

The invention relates to plate-shaped pressed bodies (wafers) ^{prepared from} ~~on the basis of~~ an inorganic sorbent and a binder, having a thickness of less than 700 μm , which are characterized by a high mechanical strength and a low brittleness and which are capable of effectively absorbing inorganic and organic gases or vapors.

Background of Invention

The manufacture of pressed bodies, in particular of tablets, on the basis of zeolites and binders is known. Thus, according to JP-A-61 15 5216, zeolite tablets are prepared by mixing a

zeolite, a binder and a lubricant, and extruding the mixture. These are ~~obviously~~ tablets with the same dimensions in all directions.

From JP-A- 56 06 3818 the manufacture of zeolite tablets for use as gas adsorbents is known. Powdered and dried (at 105 to 110°C) zeolite is mixed with 8.1% by weight of bentonite powder and ^{the mixture} is kneaded with a 4% aqueous urea solution. The mixture is ^{then} tabletted, dried and calcined at 510°C. The increase of the compressive strength ^{of the tablet} is caused by the urea content.

From JP-A-55 16 5144 it is known to knead zeolite powder for cooling aggregates in powder form with bentonite and water. ~~to extrude~~ ^{is extruded} the mixture and to form round particles with a diameter of 0.8 to 10 μ m.

According to JP-A-55 10 4913, zeolite in the Na-form is mixed with 25% by weight clay, kneaded with water, extruded, calcined at 650°C, immersed in a calcium chloride solution, washed, dried at 110°C and activated at 400°C. The tablets are used as drying agents.

According to JP-A-46 03 2572, zeolite powder is mixed with kaolin and Na- (or NH₄-)-hydroxyethyl cellulose, shaped, dried and calcined at 650°C, in order to increase the strength of the zeolite tablets.

According to JP-A-21 44 121, deodorants are obtained by extruding zeolite powder or granules with calcium chloride or bentonite and water. ~~whereafter~~ ^{then} the mixture is tabletted and ~~the tablets are~~ calcined.

According to JP-A-63 21 8234, drying agents are prepared by extruding a mixture of microporous particles (e.g. gypsum, cement, ceramic powder) and an inorganic or organic filler,

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such as CaCl_2 , LiCl , Bentonite, zeolites, PVA or other water-soluble polymers. The mixture is tabletted and subsequently hardened.

According to JP-A-60 132 643, zeolite tablets are prepared as drying agents using 20% sepiolite^e as a^a binder. The mixture is kneaded with water, tabletted, dried at 150°C and calcined at 550°C . The tablets have an improved drying efficiency as compared to bentonite tablets.

The prior art tablets are unsuitable for the ^{utilization} ~~the application~~ in a narrow space and under mechanical stresses, because they are too thick and too heavy and have ^a ~~a~~ too low ^{capacity} ~~absorption power~~ for ~~noxious~~ gases and vapors ^{based on} ~~(related to~~ mass and surface). With the methods and mixtures according to the prior art, ~~too~~ brittle pressed bodies ^{were} ~~are~~ obtained which crumble, in particular after calcining.

It is known that electroluminescent devices function ~~without~~ ~~problem~~ over a longer period of time only if a drying agent is present. ~~Such drying agents are required because of sensitivity~~ ~~this is due to the sensitivity~~ of the electrodes, in particular ~~of~~ the cathodes, ~~e.g. with respect to moisture~~ ^{generally} (cathodes ^{in use} consist of Ca or Mg alloys). Therefore, these devices are sealed as efficiently as possible under a protective gas.

The use of a moisture absorber in an electroluminescent device is disclosed in EP 500 382 A2. The ^{moisture absorber is} ~~drying agent~~ in the form of a powder or small spheres ~~is~~ applied to a black silicone resin coating. According to the preferred embodiment, the drying agent is contained in a gas-permeable bag.

Likewise, US-A-5,882,761 discloses the use of a drying agent in an electroluminescent device. The preferred drying agent is BaO . See also U.S. Patent No. 5,591,379 which discloses a coating for use with microelectronic devices comprising a desiccant powder blended with a binder.

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easily colored by the addition of pigments during manufacture.

The pressed bodies according to the invention may be produced in automatic processes in large numbers per unit time. They may be easily handled and may, for instance, be removed from a storage container by so called "pick-and-place" machines, and ~~may be~~ ^{then} inserted in an electronic device.

The pressed bodies according to the invention are capable of absorbing ~~other~~ gases (ammonia, amines, oxygen) in addition to water vapor. Since they have a high sorption capacity, the electronic device into which they are to be inserted does not have to be completely sealed air-tight, i.e. the diffusion rate for water vapor into the device can be greater than zero. Moreover, the choice of a suitable material for sealing the device (e.g. an epoxy resin), is facilitated, as the critical time in which the said material used must have reached its final and lowest permeability for water vapor can be prolonged by the use of the wafer.

Preferably, the inorganic sorbent is a natural or synthetic zeolite. However, other sorbents, such as amorphous silica or aluminium hydroxide, or mixtures of two or more sorbents, may also be used.

In general, any binder which is ~~deemed suitable~~ ^{known to} by a person skilled in the art may be used ^{the} as binder. Preferably, a ~~smectite~~ ^{smectite} clay, especially bentonite, ~~will be~~ ^{is} used ^{the} as binder. Moreover, the use of other inorganic binders, ^{such} as aluminium oxide-hydroxide (pseudo-Boehmite), is ~~feasible~~ ^{permissible}. However, ⁰ organic binders based on carbohydrates or proteins may also be used, for instance starch, cellulose derivatives (such as CMC or CEC), casein, ^{as well as} ~~but also~~ synthetic polymers, such as PVA, PVP or polyphenols or tannin-containing binders (quebracho). Also, mixtures of different binders may be used.

By the addition of the bentonite to the zeolite, the sorption capacity of the latter is surprisingly not reduced. In fact, a synergistic effect can be observed, i.e. the sorption of water vapor by the mixture is far less reduced ^{than would} ~~as could~~ be expected by ^{mere} ~~pure~~ calculation, based on the percentages of the bentonite added to the composition.

The thickness of the wafers is preferably about 200 to 400 μm , its binder content is preferably about 40 to 50% by weight.

The invention also relates to a process for preparing the pressed bodies as defined above, which process is characterized in that a mixture comprised of or containing at least one inorganic sorbent, at least one binder and optionally water and compressing aids is compressed at a pressure of at least about 70 MPa, wherein in the mixture the weight ratio of dry sorbent to dry binder is between about 4 and 0.7 and the water content of the mixture, measured at 160°C, is between about 8 and 20%; and calcining the resulting ~~green~~ pressed bodies at temperatures of at least about 500°C until the water content is ^{substantially} ~~largely~~ removed.

It has been found that by this process results in particularly advantageous pressed bodies with excellent physical and chemical properties. Especially advantageous pressed bodies are obtained at a mixing ratio of dry sorbent to binder in the initial mixture between about 1.5 and 1.

The desired water content of the mixture may be adjusted via the water content of the components (sorbent, binder) and/or ^{by} addition of water.

The A-zeolite preferably used as a sorbent is available in powder form and has a moisture content of about 10 to 22%. The bentonite preferably used as ^a binder is available in powder form with a water content of about 10 to 20. The bentonite

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used has a montmorillonite content of preferably more than 80%, based on ~~the~~^{its} dry ~~state~~^{weight}. The preferred compression aids are fatty-acid salts of a divalent or trivalent metal, such as calcium, magnesium or aluminium stearate.

It has been found that the best results may be obtained ~~in~~^{during} compression of the mixture to pressed bodies if the mixture does not comprise larger amounts, i.e. not more than about 15%, preferably not more than about 8%, in particular 0% of particles > 250 μm , preferably > 200 μm , and in particular > 150 μm , and if the majority of the particles, i.e. at least 50%, preferably at least 60%, is larger than about 45 μm .

According to a preferred process ~~feature~~, the zeolite and bentonite powders are mixed in the desired ratio with a sufficient amount of water to allow granulation of the mixture. Preferably, an intensive mixer is used herein. The amount of water which ~~has to be~~^{is} added depends on the mixing ratio of zeolite and bentonite as well as the colloid chemical properties of the bentonite used and may be easily determined by an artisan. After granulation, the mixture is adjusted or dried, respectively, to a water content of about 8 to 20%, the water content being determined at 160°C. Subsequently, the mixture is comminuted to particle sizes < 250 μm , preferably < 200 μm , in particular < 150 μm as indicated above.

It has been surprisingly found that when compressing the mixture to obtain ~~green~~ pressed bodies, the best results may be obtained if the majority of the particles of the mixture have a more or less spherical form as may be obtained e.g. by spray drying. Therefore, according to a particularly preferred embodiment, zeolite and bentonite powder ~~is~~^{are} slurried in water using a high-shear stirring device, such as a Ultra Turrax stirrer in order to obtain a pumpable slurry which is then spray-dried by conventional methods. The water content of the mixture may be adjusted to the preferred ~~values~~^{range} between about

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8 and 20% (the water content is determined at 160°C) by controlling the spray-drying process. The adjustment of the particle size distribution so that the mixture, as defined above, does not contain large~~x~~ amounts of particles $> 250 \mu\text{m}$, preferably $> 200 \mu\text{m}$, and in particular $> 150 \mu\text{m}$, and that the majority of the particles is larger than about $45 \mu\text{m}$, may also be achieved by appropriate control of the spray-drying process and optionally subsequent deagglomeration, screening and sorting steps as known in the art.

The forming of the pressed bodies from the mixture is performed at a pressure of at least about 70 MPa. The preferred pressure is between 100 and 1300 MPa. The compression of the mixture can be performed in commercially available automatic presses, the construction of which is known to the artisan.

In order to allow an efficient production of the pressed bodies, it is important that the formed bodies ~~are~~ separated easily and without residual matter from the compression tools. This can be achieved by appropriate choice of surface-modified tools (e.g. steel modified with TiN- or WC), and by ensuring precise control and ^{adjustment} ~~adjustion~~ of the water content of the mixture and control of temperature and humidity at the site of compression. At low water contents, ~~of the mixture~~ it is advantageous to ^{provide} ~~adjust~~ a relatively high humidity, e.g. 60 to 80% relative humidity at about 25 to 35°C. At high water contents, ~~of the mixture~~ a relatively low absolute humidity is preferred, e.g. 30 to 50% relative humidity at about 20 to 30°C. According to another preferred embodiment anti-sticking agents, such as magnesium or calcium stearate, are applied to the pressure tools after a predetermined number of compression cycles, e.g. after each or each second cycle. Thus, the sticking of the pressed bodies to the compression tools can be effectively avoided.

The pressed bodies are calcined at about 500 to 900°C, preferably at about 650°C, until a constant weight is obtained and the water content is ^{substantially} ~~largely removed~~.

Furthermore, it has been surprisingly found that the arching and bulging of the pressed bodies during calcination may be practically eliminated by the application of pressure to the pressed bodies during the calcination step.

According to a preferred embodiment, the application of pressure to the pressed bodies during calcination is provided by the use of a specially designed belt calcination device wherein the application of pressure to the pressed bodies is achieved via belts. In general, the application of pressure to the pressed bodies during calcination can be achieved by any method, as long as the pressure applied ~~does on the one hand~~ effectively ^{suppresses} ~~suppress~~ the bulging (curving) of the pressed bodies during calcination and ~~on the other hand the pressure~~ does not damage the pressed bodies. In general, a pressure between 10 and 30 000 Pa, in particular between 100 and 5.000 Pa, may be applied. ^{It} According to a further preferred embodiment, a certain ^{quantity} ~~amount~~ of pressed bodies is stacked in tubes, e.g. tubes made of stainless steel or ceramics. Preferably, the tubes are provided with openings (drilling holes) allowing for the evaporation of water during calcination. This ensures a rapid and uniform drying. The whole stack within a tube is subjected to a pressure sufficient to suppress bulging of the pressed bodies during calcination ^{without causing} ~~but does not cause~~ fracture, sticking or sintering of the pressed bodies during this process step. In general, the pressure lies between about 10 and 30.000 Pa, ^{preferably} ~~in particular~~ between 100 and 5,000 Pa. The wafers calcined under application of pressure according to the invention, are flat and show no or only a minimal curvature or bulging, which is a prerequisite for ~~the~~ use in electroluminescent devices.

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According to a further preferred embodiment the calcination temperature is increased stepwise in order to prevent the formation of fractures and cracks in the pressed bodies due to fast, non-uniform evaporation of water, ~~respectively~~.

The pressed bodies may also be calcined under vacuum and cooled, whereby they are capable of sorbing permanent gases, such as oxygen.

Furthermore, the pressed bodies may contain coloring pigments, such as Fe_3O_4 .

The invention further relates to the use of the above-defined pressed bodies as inserts in electronic devices or elements, such as displays, in particular electroluminescent elements, such as organic light-emitting diodes (LED). However, they may also be used in moisture-sensitive liquid crystal displays (LCD).

These devices or elements may ~~also~~ be damaged by inorganic or organic gases or vapors during manufacture or during use. ^{However,} in view of their construction, they have only limited space available for a sorption agent.

These electronic devices or elements (e.g. displays in motor vehicles and mobile telephones) are frequently exposed to strong shocks. ^{It} is therefore important that the pressed bodies ~~do~~ not break or crumble. In view of their strength, it is not necessary to cover the pressed bodies with a gas-permeable foil, which simplifies the manufacture of the electronic elements.

As compared to BaO , a significant volume and cost reduction of the electronic element may be achieved. Thus, the wafers (related to their mass) have a higher sorption capacity and sorption rate for water vapor within the required temperature

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and moisture range in an electronic element. In addition, when using BaO, it should be noted that an increase of the volume of the material by 100% occurs during the hydration reaction. Therefore, an additional volume for the expansion of the drying agent ^{must} ~~is to~~ be provided within the element, and a water-vapor permeable foil ^{must be} ~~has to~~ applied between ^{the} BaO and the electroluminescent layer ~~so as~~ to avoid ~~the~~ contact between the expanding and possibly crumbling drying agent and the layer. In contrast, the wafers (according to the invention) do not show a change of volume when absorbing water, and remain mechanically stable. Therefore, the ^{presence} ~~provision~~ of an additional expansion volume with the element and the application of a protective foil ^{are} ~~is~~ not necessary.

BaO has the additional disadvantage that it or its hydration products ^{exhibit} ~~show~~ a strong basic reaction. Furthermore, there is a strong local heating when absorbing moisture. In direct contact with organic compounds it ^{also} ~~has~~ a tendency toward self-ignition. This limits the selection of polymers for the above-mentioned protecting foils to very expensive polymers, e.g. fluoropolymers, and thus increases the costs of the element. In addition, when using BaO, disposal problems arise because, being a hazardous chemical, the dismantling, re-use and disposal of the individual parts of the electronic elements are rendered difficult.

The pressed bodies according to the invention can also be used for other purposes, e.g. as inserts for pharmaceutical packages, because ^{of the} ~~in this case only a~~ limited volume ^{which} ~~is~~ available ^{within the package} ~~for the accomodation~~ of a drying agent.

The pressed bodies may be present in any form, for instance they may be round, square, triangular or rectangular, and ~~they~~ may include holes and/or voids. The pressed bodies according to the invention are dust-free and abrasion-resistant. They

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may be manufactured in common press automats in a very large number per unit time.

The invention is illustrated by the following examples:

Example 1 (comparison)

75.2 kg zeolite 4A (water content 20%), 23.8 kg bentonite (water content 12%) and 1 kg calcium stearate ^{were} ~~are~~ mixed in an intensive mixer for 2 minutes. ~~Then,~~ ^{water} ~~was~~ added until there ^{was} ~~is~~ a ^{substantial} ~~strong~~ increase ⁱⁿ ~~of~~ viscosity, and mixing ^{was} ~~is~~ continued for 4 minutes. The mixture ^{was} ~~is~~ dried to a water content of 12% at 110°C, and subsequently granulated (Stokes granulator) and sieved (250 μ m). 0.22 g of the material with a particle size of < 250 μ m ^{were} ~~are~~ pressed to a round wafer under a pressure of 69 MPa. The ~~green~~ ^{were} ~~are~~ wafers ^{were} ~~are~~ calcined at 650°C for 3 hours, cooled with the ^{exclusion} ~~excursion~~ of moisture and packed airtight. The thickness of the wafers ^{increased} ~~increases~~ by about 15 to 25% upon calcination.

Product properties:

Thickness:	300 \pm 50 μ m
Moisture content (after calcination):	< 1%
Production rejects:	> 90%
Dropping test*:	100% fracture, wafer crumbles at the periphery

* The measure for the compression strength is the so called dropping test, wherein 100 calcined pressed bodies (round discs with a diameter of 27 μ m) are dropped from a height of 1 m with the flat side facing downwards. The percentage of the

broken test bodies ^{was} ~~is~~ determined.

Example 2 (comparison)

57 kg zeolite 4A (water content 20%), 42 kg bentonite (water content 12%) and 1 kg calcium stearate ^{were} ~~are~~ mixed in an intensive mixer for 2 minutes. ~~Then,~~ ^{was} water ^{was} ~~is~~ added until there ^{was} ~~is~~ a ^{substantial} ~~strong~~ increase of viscosity, and mixing ^{was} ~~is~~ continued for 4 minutes. The mixture ^{was} ~~is~~ dried to a water content of 12% at 110°C, and subsequently granulated (Stokes granulator) and sieved (250 μ m). 0.22 g of the material with a particle size of < 250 μ m ^{were} ~~are~~ pressed to a wafer under a pressure of 69 MPa. The ~~green~~ ^{were} ~~are~~ wafers ^{were} ~~are~~ calcined at 650°C for 3 hours, cooled with the exclusion of moisture and packed airtight.

Product properties:

Thickness:	300 \pm 50 μ m
Moisture content (after calcination):	< 1%
Production rejects:	75%
Dropping test:	80% fracture

Example 3

57 kg zeolite 4A (water content 20%), 42 kg bentonite (water content 12%) and 1 kg calcium stearate ^{were} ~~are~~ mixed in an intensive mixer for 2 minutes. ~~Then,~~ ^{was} water ^{was} ~~is~~ added until there is a strong increase of viscosity, and mixing ^{was} ~~is~~ continued for 4 minutes. The mixture ^{was} ~~is~~ dried to a water content of 12% at 110°C, and subsequently granulated (Stokes granulator) and sieved (250 μ m). 0.22 g of the material with a

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particle size of $< 250 \mu\text{m}$ ^{were} ~~are~~ pressed to a wafer under a pressure of 72 MPa. The green wafers ^{were} ~~are~~ treated as in Example 2.

Product properties:

Thickness:	$300 \pm 50 \mu\text{m}$
Moisture content (after calcination):	$< 1\%$
Production rejects:	$< 50\%$
Dropping test:	60% fracture

Example 4

57 kg zeolite 4A (water content 20%), 42 kg bentonite (water content 12%) and 1 kg calcium stearate ^{were} ~~are~~ mixed in an intensive mixer for 2 minutes. Then, ^W ~~water~~ ^{was} added until there is a ^{substantial} ~~strong~~ increase of viscosity, and mixing ^{was} ~~is~~ continued for 4 minutes. The mixture ^{was} ~~is~~ dried to a water content of 12% at 110°C , and subsequently granulated (Stokes granulator) and sieved ($250 \mu\text{m}$). 0.22 g of the material with a particle size of $< 250 \mu\text{m}$ ^{were} ~~are~~ pressed to a wafer under a pressure of 350 MPa. The ~~green~~ wafers ^{were} ~~are~~ calcined at 650°C for 3 hours, cooled with the exclusion of moisture and packed air-tight.

Product properties:

Thickness:	300 ± 50 μm
Moisture content (after calcination):	< 1%
Production rejects:	< 25%
Dropping test:	15% fracture
Sorption capacity*	
after 1 hour	5.4% by weight
after 5 hours	7.2% by weight
after 24 hours	13.0% by weight

* The sorption capacity for water vapor ^{was} ~~is~~ determined at 25°C in an atmosphere having a moisture content of 10%.

Example 5

The procedure of Example 4 was repeated except that the calcination of the ~~green~~ wafers was carried out in vacuum. The calcined wafers had essentially the same product properties as the wafers of Example 4, but they additionally showed a sorption capacity for oxygen of about 5 ml/g (determined in a dry oxygen atmosphere).

Example 6

56.5 kg zeolite 4A (water content 20%), 41.5 kg bentonite (water content 12%), 1 kg calcium stearate and 1 kg quebracho ~~were~~ ^{were} mixed in an intensive mixer for 2 minutes. ~~Then, water is~~ ^{was} added until there ~~is~~ ^{was} a ~~strong~~ ^{substantial} increase of viscosity, and mixing ~~is~~ ^{was} continued for 4 minutes. The mixture ~~is~~ ^{was} dried to a water content of 12% at 110°C, and subsequently granulated (Stokes granulator) and sieved (250 μm). 0.22 g of the

material with a particle size of $< 250 \mu\text{m}$ ^{were} ~~are~~ pressed to a wafer under a pressure of 200 MPa. The ~~green~~ ^{were} wafers ~~are~~ calcined at 650°C for 3 hours, cooled with the exclusion of moisture and packed air-tight.

Product properties:

Thickness:	$300 \pm 50 \mu\text{m}$
Moisture content (after calcination):	$< 1\%$
Production rejects:	$< 35\%$
Dropping test:	10% fracture

Example 7

The procedure of Example 4 was repeated except that the compression pressure was 1200 MPa. The dropping test yielded 10% fractures. The rejects were $< 10\%$.

Example 8

The procedure of Example 4 was repeated except that 54 kg zeolite 4A, 40 kg bentonite, 5 kg Fe_3O_4 and 1 kg calcium stearate are used. The resulting wafers were dark in color and could be used for a contrast surface in a LED display.

Example 9

110.5 kg zeolite 4A (water content 20%), 76.0 kg bentonite (water content 12%) and 1.9 kg calcium stearate ^{were} ~~are~~ slurried in a high-shear stirring device in sufficient water to obtain a pumpable slurry. The slurry (suspension) ^{was} ~~is~~ spray-dried so

that a powder with a water content of 9.2% (determined by drying at 160°C) and a particle size distribution of 0% > 150 μm and 30% < 45 μm ^{was} ~~is~~ obtained. 0.17 g of this material are compressed to a wafer having a diameter of 20 mm using a pressure of 190 MPa, the partial pressure of water vapor in the environment (air) surrounding the compression tools set to about 30 mbar. The ~~green~~ ^{were} wafers ~~are~~ calcined under the application of pressure for 3 hours at 650°C. ~~Thus, 300 green~~ wafers ~~each are~~ ^{were} stacked in ~~stainless~~ ^{stainless} steel tubes with drilling holes (openings) (height 120 mm, inner diameter 20 mm), and the stack ^{was} ~~is~~ subjected to a pressure of 2.550 Pa. Following calcination, the wafers ^{were} ~~are~~ cooled under exclusion of moisture and packed.

Product properties:

Thickness:	300 \pm 50 μm
Vertical dimension: (thickness + curvature (bulging))	< 350 μm
Moisture content (after calcination):	< 1%
Production rejects:	< 5%
Dropping test:	< 1%

Example 10

110.5 kg zeolite 4A (water content 20%), 76.0 kg bentonite (water content 12%) and 1.9 kg calcium stearate ^{were} ~~are~~ slurried in a high-shear stirring device (Ultra-Turrax stirrer) in sufficient water to obtain a pumpable slurry. The slurry (suspension) ^{was} ~~is~~ spray-dried so that a powder with a water content of 12.6% (determined by drying at 160°C) and a particle size distribution of 0% > 150 μm and 26% < 45 μm ^{was} ~~is~~ obtained. 0.17 g of this material ^{was} ~~are~~ compressed to a wafer

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having a diameter of 20 mm using a pressure of 210 MPa, the partial pressure of water vapor in the environment (air) surrounding the compression tools ^{was} being about 17 mbar. The ~~green~~ ^{were} wafers ~~are~~ calcined under the application of pressure for 3 hours at 650°C. Thus, 300 ~~green~~ wafers ~~each are~~ ^{were} stacked in stainless steel tubes with drilling holes (openings) (height 120 mm, inner diameter 22 mm), and the stack ~~is~~ ^{was} subjected to a pressure of 2.550 Pa. After calcination, the wafers ~~are~~ ^{were} cooled under exclusion of moisture and packed.

Product properties:

Thickness: 300 ± 50 μm
Vertical dimension: < 350 μm
(thickness + curvature (bulging))
Moisture content
(after calcination): < 1%
Production rejects: < 5%
Dropping test: < 1%

Example 11

110.5 kg zeolite 4A (water content 20%), 76.0 kg bentonite (water content 12%) and 1.9 kg calcium stearate ^{were} ~~are~~ slurried in a high-shear stirring device (Ultra-Turrax stirrer) in sufficient water to obtain a pumpable slurry. The slurry (suspension) ^{was} ~~is~~ spray-dried so that a powder with a water content of 15.5% (determined by drying at 160°C) and a particle size distribution of 4% > 150 μm and 8% < 45 μm ^{was} ~~is~~ obtained. 0.17 g of this material ^{were} ~~are~~ compressed to a wafer having a diameter of 20 mm using a pressure of 195 MPa, the partial pressure of water vapor in the environment (air) surrounding the compression tools ^{was} ~~is~~ set to about 12 mbar. The ~~green~~ ^{were} wafers ~~are~~ calcined under the application of pressure

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for 3 hours at 650°C. Thus, 300 green wafers ^{were} ~~each are~~ stacked in stainless steel tubes with drilling holes (openings) (height 120 mm, inner diameter 22 mm), and the stack ^{was} ~~is~~ subjected to a pressure of 2.550 Pa. After calcination, the wafers ^{were} ~~are~~ cooled under exclusion of moisture and packed.

Product properties:

Thickness: 300 ± 50 μm
Vertical dimension: < 350 μm
(thickness + curvature (bulging))
Moisture content
(after calcination): < 1%
Production rejects: < 5%
Dropping test: < 1%

Example 12

57 kg zeolite 4A (water content 20%), 42 kg of a 50/50 mixture of attapulgite and kaolin (water content 12%) and 1 kg calcium stearate were mixed for 2 minutes in an intensive mixer. Then, water ^{was} ~~is~~ added until there ^{was a substantial} ~~is a strong~~ increase of viscosity, and mixing ^{was} ~~is~~ continued for 4 minutes. The mixture ^{was} ~~is~~ dried to a water content of 12%, and subsequently granulated and sieved (150 μm sieve). 0.17 g of this material having a particle size < 150 μm ^{were} ~~are~~ pressed to a wafer under a pressure of 200 MPa. The ~~green~~ wafers ^{were} ~~are~~ calcined for 3 hours at 650°C, cooled with the exclusion of moisture and packed.

Product properties:

Thickness:	300 ± 50 μm
Moisture content (after calcination):	< 1%
Production rejects:	25%
Dropping test:	70% fracture

Example 13

An organic electroluminescent element(1) (square, area 12.9 cm²), as shown in Fig. 1, ^{was} ~~is~~ produced using a wafer (circular, diameter 27 μm) of Example 4. After fixing the wafer(2) to the back wall(3) of the element, it ^{was} ~~is~~ adhered to the glass substrate(5) of the element by means of an adhesive(4) and it ~~was~~ sealed as far as possible by means of an adhesive. Then, a microphotograph (enlarged 50 times) of the light-emitting part (6) (consisting of the anode(7), the light-emitting layer(8) and the cathode(9) of the element ^{was} ~~is~~ taken. This photograph does not show dark (non-luminescent) spots, which ~~would~~ indicate an attack on cathode(9).

The element ^{was} ~~is~~ exposed to a temperature of 85°C and a relative humidity of 85% for 500 hours. Subsequently, a new microphotograph of the light-emitting part(6) of element(1) ^{was} ~~is~~ taken. A comparison of the two photographs shows that no dark spots were formed, which ~~would~~ indicate an attack on the cathode(9).

Example 14 (comparison)

An organic electroluminescent element(1) as in Example 9 ^{was}~~is~~ produced using BaO. As a cover for the BaO, a water-permeable teflon foil ^{was}~~is~~ used which ^{was}~~is~~ attached to the back wall(3) by means of a thin double-adhesive tape. The amount of BaO ^{was}~~is~~ adjusted so that the total mass of BaO, of the teflon foil and the double-adhesive tape exactly corresponds to that of the wafer produced in Example 9. Thereafter, as described in Example 9, enlarged photographs of the light-emitting element before and after a storage at 85°C and 85% humidity for 500 hours ^{were}~~are~~ taken. A comparison of the two photographs shows ^{ed}~~a~~ significantly recognizable growth of dark spots which indicated an attack on cathode(9).

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Claims

1. Plate-shaped pressed body (wafer), comprising at least one inorganic sorbent and at least one binder, having a thickness of less than 700 μm , which is obtainable by compressing a mixture, comprised of or containing, respectively, the inorganic sorbent(s) and the binder(s) at a pressure of at least 70 MPa, wherein in the mixture the weight ratio of dry sorbent(s) to dry binder(s) is between about 4 and 0.7 and the water content of the mixture, determined at 160°C, is between about 8 and 20%; and calcining the resulting green pressed bodies at temperatures of at least 500°C, until the water content is largely removed.
2. Pressed body according to claim 1, characterized in that water and/or compression aids such as fatty-acid salts of a divalent or trivalent metal are added to the mixture.
3. Pressed body according to claims 1 or 2, characterized in that the calcination is continued to constant weight or to a residual moisture of < 2% by weight, measured at the calcination temperature.
4. Pressed body according to any one of claims 1 to 3, characterized in that the inorganic sorbent is a natural or synthetic zeolite.
5. Pressed body according to any one of claims 1 to 4, characterized in that the binder is a smectitic clay, preferably bentonite.

6. Pressed body according to any one of claims 1 to 5, characterized in that the thickness of the wafer is about 200 to 400 μm .

7. Pressed body according to any one of claims 1 to 6, characterized in that in the mixture the weight ratio of dry sorbent to dry binder is between about 1,5 and 1.

8. Pressed body according to any one of claims 1 to 7, characterized in that the pressure is about 100 to 1300 MPa.

9. Pressed body according to any one of claims 1 to 8, characterized in that it has been compressed with a compression aid, preferably a fatty acid salt of a divalent or trivalent metal.

10. Pressed body according to any one of claims 1 to 9, characterized in that the mixture does not comprise large amounts, preferably not more than 15%, more preferred not more than 8% and particularly preferred 0% of particles $> 250 \mu\text{m}$, preferably $> 200 \mu\text{m}$ and particularly preferred $> 150 \mu\text{m}$.

11. Pressed body according to any one of claims 1 to 10, characterized in that the majority of particles in the mixture, e.g. at least about 50%, in particular at least about 60%, is larger than about 45 μm .

12. Pressed body according to any one of claims 1 to 11, characterized in that the mixture mainly comprises spherical particles.

13. Pressed body according to any one of claims 1 to 12, characterized in that more than 50%, preferably more than 75%, more preferably more than 80% and most preferably more than 98% of the particles are essentially spherical.

14. Pressed body according to any one of claims 1 to 13, characterized in that the mixture is obtained by spray drying.

15. Pressed body according to any one of claims 1 to 14, characterized in that it has been calcined under vacuum.

16. Pressed body according to any one of claims 1 to 15, characterized in that the calcination is performed with application of pressure to the pressed bodies.

17. Pressed body according to claim 16, characterized in that the calcination of the pressed bodies under pressure is performed in a perforated tube.

18. Process for producing plate-shaped pressed bodies (wafers), characterized in that a mixture comprised of or containing, respectively, inorganic sorbent(s) and binder(s) is compressed at a pressure of at least 70 MPa, wherein in the mixture the weight ratio of dry sorbent(s) and dry binder(s) is between about 4 and 0.7 and the water content of the mixture, measured at 160°C, is between about 8 and 20%; and calcining the resulting green pressed bodies at temperatures of at least about 500°C until the water content is largely removed.

19. Process according to claim 19, characterized in that one or more of the components referred to in any one of claims 2 to 17 are used under the conditions recited therein.

20. The use of the pressed body according to any one of claims 1 to 17 as insert in electronic devices, such as displays, in particular in electroluminescent elements.

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Abstract
Summary

There is disclosed a plate-shaped pressed body (wafer) ^{produced} ~~on the~~
~~from~~ ~~basis of~~ an inorganic sorbent and a binder, having a thickness
of less than 700 μm , which is ^{produced by the process of} ~~obtainable by~~ compressing a
mixture of the inorganic sorbent, the binder, and optionally
water and compression aids at a pressure of at least 70 MPa,
wherein ~~in the mixture~~ the weight ratio of dry sorbent to dry
binder is between about .4 and 0.7 and the water content of the
mixture, measured at 160°C, is between 8 and 20%; and
calcining the resulting ~~green~~ pressed body at temperatures of
at least about 500°C, until the water content is ^{substantially} ~~largely~~
removed.

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